IN THE UNITED STATES PATENT AND TRADEMARK OFFICE

I, SALLY LESLEY HEDLEY, B.A., M.I.L., M.I.T.I., declare

- That I am a citizen of the United Kingdom of Great Britain and Northern Ireland, residing at 29 Parkholme Road, London E8 3AG.
- 2. That I am well acquainted with the German and English languages.
- 3. That the attached is a true translation into the English language of the Request and Specification as originally filed of International Patent Application No. PCT/EP99/09240, save that the English version of the title has been taken from the cover page of the corresponding International Publication No. WO 00/34388.
- 4. That all statements made herein of my own knowledge are true and that all statements made on information and belief are believed to be true; and further that these statements are made with the knowledge that wilful false statements and the like so made are punishable by fine or imprisonment, or both, under Section 1001 of Title 18 of the United States Code and that such wilful false statements may jeopardise the validity of the patent application in the United States of America or any patent issuing thereon.

DECLARED THIS 10 to DAY OF MAY 2001

SALLY HEDLEY

Sally Hedley.

PCT



REQUEST

The undersigned requests that the present international application be processed according to the Patent Cooperation Treaty.

For	receiving	Office	use	only	
-----	-----------	--------	-----	------	--

PCT/EP 99/0924Q

International Application No.

29 NOV 1999

(29.11.99)

International Filing Date

EUROPEAN PATENT OFFICE PCT INTERNATIONAL APPLICATION

	Name of receiving Office and "F	CI International Application
	Applicant's or agent's file refere (if desired) (12 characters maxir	num) HE/K-21888/A
ox No. I TITLE OF INVENTION		
Hydrophobic epoxide resin sy	rstem	
ox No. II APPLICANT		
ame and address: (Family name followed by given name; for a legal entit oust include postal code and name of country. The country of the address i tate (that is, country) of residence if no State of residence is indicated bel	indicated in this Box is the applicant 3	This person is also inventor
Ciba Specialty Chemicals Hol	lding Inc.	Telephone No. +41 61 636 11 11
Klybeckstrasse 141		
4057 Basle		Facsimile No. +41 61 636 79 76
СН		Teleprinter No.
State (that is, country) of nationality: CH	State (that is, country) of residence	СН
	mica builds except	United States the States indicated in the Supplemental Box
Box No. III FURTHER APPLICANT(S) AND/OR	(FURTHER) INVENTOR(S)	
Name and address. (Full the Joint of State of February) of the address State (that is, country) of residence if no State of residence is indicated be BEISELE, Christian Lindenweg 36 79424 Auggen DE	State (that is, country) of residence	This person is: applicant only applicant and inventor inventor only (If this check-box is marked, do not fill in below.)
State (that is, country) of nationality:	State (mar is, country) or resident	DE
		the States indicated in the Supplemental Box
Further applicants and/or (further) inventors are	indicated on a continuation sheet	•
Box No. IV AGENT OR COMMON REPRESEN	TATIVE; OR ADDRESS FOR	CORRESPONDENCE
The person identified below is hereby/has been appointed to a of the applicant(s) before the competent International Authorit	ct on behalf agent ites as:	common representative
Name and address: (Family name followed by given name; for a le address must include postal code and name of	egal entity, full official designation. The country.)	Telephone No. +41 61 636 11 11
Ciba Specialty Chemicals Ho Patent Department Klybeckstrasse 141	lding Inc.	Facsimile No. +41 61 636 79 76
4057 Basle		Teleprinter No.

Form PCT/RO/101 (first sheet) (July 1999)

See Notes to the request form

RAINMOILER, Thomas Schauinslandstrasse 2 79576 Weil am Rhein DE State (that is, country) of nationality: AT State (that is, country) of residence: DE State (that is, country) of residence: States This person is applicant and inventor inventor only (If this check-b is marked, do not fill in below the Supplemental Events and address; family name followed by given name, for a kept entity, full official distipation. The address must be supplemental to the Supplemental Events of America only of residence is indicated below.) TANG, Qian State (that is, country) of residence is indicated below.) TANG, Qian State (that is, country) of residence is indicated below.) State (that is, country) of residence only (If this check-lis marked, do not fill in below the Supplemental Country) of residence of a supplicant only (If this check-lis marked, do not fill in below the supplemental Country) of residence of the United States of America only of the States indicated below.) State (that is, country) of residence is indicated below.) State (that is, country) of residence of the Supplemental Country of the States indicated in this Bos is the applicant. The address must be supplicant only (If this check-lis marked, do not fill in below the supplemental Country) of residence. CH This person is applicant only applicant only of the States indicated below.) State (that is, country) of residence is indicated below.) State (that is, country) of residence is indicated below.) State (that is, country) of residence is indicated below.) This person is applicant only in the States indicated below.) State (that is, country) of residence is the Supplemental Country of residence is indicated below.) State (that is, country) of residence is the States indicated in this Bos is the applicant. The address must be supplicant only inventor only (If this check-lis marked, do not fill in below the Supplemental in the supplicant only inventor only (If this check-lis marked, do not fill in below the supplicant only of America only of	Continuation of Box No. III FURTHER APPLICANT(S) AND/OR (FURTHER) IN	NTOR(S)
This person is applicant only TANG, Qian State (that is, country) of nationality: CN CN CN This person is applicant only State (that is, country) of nationality: CN This person is applicant only (If this check-by is marked, do not fill in below the States indicated the United States of America This person is applicant This person is applicant only TANG, Qian TANG, Qian State (that is, country) of nationality: CN This person is applicant This person is applicant only applicant and inventor Inventor only (If this check-by is marked, do not fill in below TANG, Qian State (that is, country) of nationality: CN CN CH This person is applicant This person is applicant This person is applicant only applicant only applicant only applicant only applicant only This person is applicant This person is applicant This person is applicant only applicant only applicant only applicant only This person is applicant This person is applicant only applicant and inventor inventor only (If this check-by is marked, do not fill in below This person is applicant only applicant and inventor inventor only (If this check-by is marked, do not fill in below This person is applicant This person is applicant This person is applicant This person is applicant only applicant and inventor inventor only (If this check-by is marked, do not fill in below This person is applicant only applicant and inventor inventor only (If this check-by is marked, do not fill in below This person is applicant only applicant and inventor inventor only (If this check-by is marked,			d in the request.
This person is applicant State (that is, country) of residence (Family mame followed by given name: for a legal entity) TANG, Qian State (that is, country) of mationality: State (that is, country) of residence (Family mame followed by given name: for a legal entity, full official designation. The address must include postal code and name of country. The country of the address indicated in this Bot to the applicant? State (that is, applicant and inventor inventor only (If this check-to is marked, do not fill in below the supplemental in the states indicated in this Bot to the applicant? State (that is, country) of residence: CH State (that is, country) of residence (Family mame followed by given name; for a legal entity, full official designation. The address must of America only in the States indicated in this Bot is the applicant? State (that is, country) of residence (Family pame) followed by given name; for a legal entity, full official designation. The address must country) of residence (I mail pame) followed by given name; for a legal entity, full official designation. The address must country) of residence (I mail pame) followed by given name; for a legal entity, full official designation. The address must country) of residence (I mail pame) followed by given name; for a legal entity, full official designation. The address must country) of residence (I may followed by given name; for a legal entity, full official designation. The address must for the purposes of: State (that is, country) of nationality: State (that is, country) of nationality: State (that is, country) of residence is indicated below.) This person is applicant and inventor inventor only (If this check-is marked, do not fill in below the supplemental for the purposes of: State (that is, country) of residence is indicated below.) This person is applicant and inventor inventor only (If this check-is marked, do not fill in below the supplemental for the purposes of: State (that is, country) of residence is indicated below.) This	nclude postal code and name of country. The country of the address indicated in this B country) of residence if no State of residence is indicated below.) KAINMÜLLER, Thomas Schauinslandstrasse 2 79576 Weil am Rhein	l designation. The address must ox is the applicant's State (that is,	applicant only applicant and inventor
This person is applicant This person is applicant All designated and address: (Family name pollowed by given name; for a legal entity, full official designation. The address must include postal code and ame of country. The country of the address indicated in this Box is the applicant's State (that is, country) of residence: TANG, Qian State (that is, country) of nationality: CN State (that is, country) of residence: This person is applicant all designated all designated states except the United States of America This person is applicant all designated and states of the address must include postal code and name of country. The country of the address indicated in this Box is the applicant. This person is applicant This person is applicant all designated all designated states except the United States of America This person is applicant This person is applicant all designated all designated states except the United States of America This person is applicant only (if this check-tis must include postal code and name of country.) The country of the address indicated in this Box is the applicant? State (that is, country) of residence: This person is applicant This person is applicant and inventor inventor only (if this check-tis marked, do not fill in below) State (that is, country) of nationality: State (that is, country) of nationality: State (that is, country) of residence: This person is applicant and inventor inventor only (if this check-tis marked, do not fill in below) applicant and inventor inventor only (if this check-tis marked, do not fill in below) This person is applicant and inventor inventor only (if this check-tis marked, do not fill in below) This person is applicant and inventor inventor only (if this check-tis marked, do not fill in below) This person is applicant and inventor inventor only (if this check-tis marked, do not fill in below) This person is applicant and inventor inventor only (if this check-tis marked, do not fill in below) This person is applicant an	State (that is, country) of nationality: AT	State (that is, country) of reside	DE
This person is applicant all designated States except the United States of America only of residence is indicated below.) State (that is, country) of nationality: CN State (that is, country) of nationality: CH This person is applicant all designated all designated States except the United States of America only of residence: This person is applicant all designated below.) States (that is, country) of nationality: CH This person is applicant all designated below.) States (that is, country) of residence: This person is applicant all designated below.) States (that is, country) of residence is indicated below.) State (that is, country) of residence is the United States of America only of residence is the Supplemental include postal code and man of country. The country of the address indicated in this Box is the applicant is for the purposes of: This person is applicant and inventor inventor only (If this checkis marked, do not fill in below inventor only (If this checkis marked, do not fill in below inventor only (If this checkis marked, do not fill in below inventor only (If this checkis marked, do not fill in below inventor only (If this checkis marked, do not fill in below inventor only (If this checkis marked, do not fill in below inventor only (If this checkis marked, do not fill in below inventor only (If this checkis marked, do not fill in below inventor only (If this checkis marked, do not fill in below inventor only (If this checkis marked, do not fill in below inventor only (If this checkis marked, do not fill in below inventor only (If this checkis marked, do not fill in below inventor only (If this checkis marked, do not fill in below inventor only (If this checkis marked, do not fill in below inventor only (If this checkis marked, do not fill in below inventor only (If this checkis marked, do not fill in below inventor only (If this checkis marked, do not fill in below inventor only (If this checkis marked, do not fill in below inventor only (If this checkis marked, do not fill in below	for the purposes of: States the United States	of America of An	
This person is applicant all designated all designated States except the United States of America only the States indicate the Supplemental Name and address: (Family name followed by given name: for a legal entity, full official designation. The address must include postal code and name of country. The country of the address indicated in this Box is the applicant's State (that is, country) of residence if no State of residence is indicated below.) State (that is, country) of nationality: State (that is, country) of nationality: State (that is, country) of residence: This person is applicant all designated all designated States except the United States of America only the States indicated in this Box is the applicant inventor only (If this check is marked, do not fill in below the United States of America only the States indicated for the purposes of: This person is applicant all designated all designated States except the United States of America only the States indicated the Supplemental Name and address: (Family name followed by given name: for a legal entity, full official designation. The address must include postal code and name of country. The country of the address indicated in this Box is the applicant in State (that is, country) of residence is indicated below.) State (that is, country) of nationality: State (that is, country) of residence is indicated below.) Further applicant and/or (further) inventors are indicated on another continuation sheet.	include postal code and name of country. The country of the address indicated in this to country) of residence if no State of residence is indicated below.) TANG, Qian Stallenrain 1 4104 Oberwil	il designation. The address must lox is the applicant's State (that is,	applicant only applicant and inventor
This person is applicant States Sta	State (that is, country) of nationality: CN	State (that is, country) of resid	
Name and address: (Family name followed by given name; for a legal entity, full official designation. The address must include postal code and name of country. The country of the address indicated in this Box is the applicant's State (that is, country) of residence if no State of residence is indicated below.) State (that is, country) of nationality: State (that is, country) of nationality: State (that is, country) of residence: This person is applicant In United States of America only In State (that is, country) of nationality: State (that is, country) of residence: This person is applicant In United States of America only In State (that is, country) of nationality: State (that is, country) of residence: This person is applicant In United States of America only In State (that is, country) of residence is indicated below.) This person is: In person	Titls person is approved.	IV	
This person is applicant all designated all designated States except the United States the States indicate for the purposes of: the United States of America the United States the Supplemental the States indicated the Supplemental the States indicated the Supplemental the States the Supplemental the Supplemental the States the Supplemental the Supplemental the States the Supplemental the States the States the States the States the States the States the United States the United States the United States the States the Supplemental the States the Supplemental the Supplemental the States the States the Supplemental the States the Supplemental the States the Supplemental the Supplemental	Name and address: (Family name followed by given name; for a legal entity, full offici include postal code and name of country. The country of the address indicated in this country) of residence if no State of residence is indicated below.)	al designation. The address must Box is the applicant's State (that is,	applicant only applicant and inventor
This person is applicant for the purposes of: All designated states except for the purpose of: America only the Supplemental of America only in the Supplemental for the purpose of: America only the Supplemental designation. The address must for the purpose of: America only the supplemental of America only of America only of America only in the Supplemental designated states except for the purpose of: America only the Supplemental designated on another continuation sheet.	State (that is, country) of nationality:	State (that is, country) of resid	dence:
State (that is, country) of nationality: State (that is, country) of residence: State (that is, country) of nationality: State (that is, country) of nationality: This person is applicant all designated all designated for the purposes of: States and/or (further) inventors are indicated on another continuation sheet.	for the purposes of: States the United States	s of America of A	
This person is applicant all designated all designated States except the United States of America only the States indicated for the purposes of: Further applicants and/or (further) inventors are indicated on another continuation sheet.	include postal code and name of country. The country of the address indicated in this	ial designation. The address must Box is the applicant's State (that is,	applicant only applicant and inventor
for the purposes of: States all designated all designated states of America of America only the Supplementa Further applicants and/or (further) inventors are indicated on another continuation sheet.	State (that is, country) of nationality:	State (that is, country) of resi	
	Thus person is approach	tatos univer	
	Further applicants and/or (further) inventors are indicated	on another continuation sheet	.

Bo	x No.V	V DESIGNATION OF STATES					
		esignations are hereby made under 4.9(a) (mark the applicable	e check	-boxes: ai	least one (marked):		
Re	gional Paten						
×	AP	ARIPO Patent: GH Ghana, GM Gambia, KE Kenya, LS Lesotho, MW Malawi, SD Sudan, SL Sierra Leone, SZ Swaziland, TZ United Republic of Tanzania, UG Uganda, ZW Zimbabwe, and any other State which is a Contracting State of the Harare Protocol and of the PCT					
×	EA	Eurasian Patent: AM Armenia, AZ Azerbaijan, BY Belarus, KG Kyrgyzstan, KZ Kazakstan, MD Republic of Moldova, RU Russian Federation, TJ Tajikistan, TM Turkmenistan, and any other State which is a Contracting State of the Eurasian Patent Convention and of the PCT					
X	EP	European Patent: AT Austria, BE Belgium, CH and LI Switzerland and Liechtenstein, CY Cyprus, DE Germany, DK Denmark, ES Spain, FI Finland, FR France, GB United Kingdom, GR Greece, IE Ireland, IT Italy, LU Luxembourg, MC Monaco, NL Netherlands, PT Portugal, SE Sweden, and any other State which is a Contracting State of the European Patent Convention and of the PCT					
×	OA	OAPI Patent: BF Burkina Faso, BJ Benin, CF Central African Republic, CG Congo, CI Côte d'Ivoire, CM Cameroon, GA Gabon, GN Guinea, GW Guinea-Bissau, ML Mali, MR Mauritania, NE Niger, SN Senegal, TD Chad, TG Togo, and any other State which is a member State of OAPI and a Contracting State of the PCT (if other kind of protection or treatment desired, specify on dotted line)					
Na	itional Pater	at (if other kind of protection or treatment desired, specify on dotte	d line):				
×	AE	United Arab Emirates	X	LR	Liberia		
Z	_	Albania	X	LS	Lesotho		
×		Armenia	X	LT	Lithuania		
×		Austria	$\overline{\mathbf{Z}}$	LU	Luxembourg		
Ø		Australia	×	LV	Latvia		
Σ		Azerbaijan	×	MD	Republic of Moldova		
K		Bosnia and Herzegovina	×	MG	Madagascar		
		Barbados.	×	MK	The former Yugoslav Republic of Macedonia		
		Bulgaria			The former regional respection of reasonable		
		Brazil	X	MN	Mongolia		
		Belarus	×	MW	Malawi		
				MX	Mexico		
		Canada	×	MA NO	Norway		
		nd LI Switzerland and Liechtenstein	×		New Zealand		
		China	1 23	NZ pr			
		Cuba		PL	Poland		
Σ		Czech Republic		PT	Portugal		
2		Germany	X	RO	Romania		
<u> </u>		Denmark	区	RU	Russian Federation		
	₫ ee	Estonia	X	SD	Sudan		
	₫ ES	Spain	X	SE	Sweden		
	₹ FI	Finland	X	\mathbf{SG}	Singapore		
D		United Kingdom	X	SI	Slovenia		
5	₫ GD	Grenada	X	SK	Slovakia		
I	₹ GE	Georgia	X	SL	Sierra Leone		
-2	⊠ GH	Ghana	X	TJ	Tajikistan		
	S GM	Gambia	M	TM	Turkmenistan		
	HR	Croatia	\mathbf{x}	TR	Turkey		
	MU HU	Hungary	X	TT	Trinidad and Tobago		
	X ID	Indonesia	×	UA	Ukraine		
		Israel	X	UG	Uganda		
	X IN	India	×	US	United States of America		
	Zi is	leeland		-55	Office States of America.		
	_	Japan	X	UZ	Uzbekistan		
	72	•	X	VN	Viet Nam		
		Kenya		YU	Yugoslavia		
	X KG	Kyrgyzstan	×		YugosiaviaSouth Africa		
1	🔀 кр	Democratic People's Republic of Korea	X	Z.A 7W	South AiricaZimbabwe		
,	⊠ KR	Republic of Korea		ZW ck-boxes r	reserved for designating States (for the purposes of a national		
	5-73	Kazakstan			have become party to the PCT after issuance of this sheet:		
1 .		Saint Lucia	pater	CR	Costa Rica		
	~ =		X	DM	Dominica		
1,	⊠ lk	Sri Lanka	X	TZ	United Republic of Tanzania		
			120	MA	Marrico		

Precautionary Designation Statement: In addition to the designations made above, the applicant also makes under Rule 4.9(b) all other designations which would be permitted under the PCT except any designation(s) indicated in the Supplemental Box as being excluded from the scope of this statement. The applicant declares that those additional designations are subject to confirmation and that any designation which is not confirmed before the expiration of 15 months from the priority date is to be regarded as withdrawn by the applicant at the expiration of that time limit. (Confirmation of a designation consists of the filing of a notice specifying that designation and the payment of the designation and confirmation fees. Confirmation must reach the receiving Office within the 15-month time limit.)

Filing Date	Number		Where earlier application	n is:	
of earlier application (day/month/year)	of earlier application	national application: country	regional application:*	international application	
tem (1) 09 December 1998 (09.12.98)	2441/98	СН	regional Office	receiving office	
tem (2)					
tem (3)					
The receiving Office is requested to proof the earlier application(s) (only if the of the present international application where the earlier application is an ARIPO Convention for the Protection of Industrial Profession	earlier application was filed n is the receiving Office) iden application, it is mandatory to	with the Office which for ntified above as item(s): _ o indicate in the Suppleme	the purposes ntal Box at least one coun		
	ARCHING AUTHORIT				
Choice of International Searching Authorically two or more International Searching Authorities of competent to carry out the international search, individual representational search, individual representational search.	are carried out by or r	equested from the Internation	al Searching Authority):	(if an earlier search has beer	
ISA/				,,, ,, ,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,	
Box No. VIII CHECK LIST; LANGU	JAGE OF FILING			·	
This international application contains the following number of sheets:	This international applica	ation is accompanied by	the item(s) marked below	:	
equest : 4	1. A fee calculation	on sheet			
escription (excluding : 21 equence listing part)					
laims : 3	4. statement explaining lack of signature				
bstract : 1	5. priority document(s) identified in Box No VI as item(s): (1)				
rawings : ——	6. translation of international application into (language):				
equence listing part f description : ——	7. separate indi	cations concerning deposit	ed microorganism or othe	er biological material	
Total number of sheets : 29	8. In nucleotide ar	nd/or amino acid sequence	listing in computer readal	ble form	
Figure of the drawings which hould accompany the abstract:		Language of filing of international application			
Box No. IX SIGNATURE OF APPI	LICANT OR AGENT				
lext to each signature, indicate the name of the request)	he person signing and the cap	acity in which the person	signs (if such capacity is r	not obvious from reading	
e request,		Ciba Specialty	Chemicals Ho	lding Inc	
		(sign	ature)		
25.11.1999		Verena Sp Patent Admi	_		
	For receiving	Office use only			
. Date of actual receipt of the purported international application:	29 NOV 1		11.99)	2. Drawings:	
 Corrected date of actual receipt due to late timely received papers or drawings complethe purported international application: 				not	
Date of timely receipt of the required				receive	
corrections under PCT Article 11(2): International Searching Authority specified by the applicant:	<i>y</i>	6. Transmittal o	f search copy delayed		

Form PCT/RO/101 (last sheet) (July 1999)

- 1 -

Hydrophobic epoxide resin system

The present invention relates to a composition comprising an epoxy resin, various polysiloxanes and fluorinated compounds, to crosslinked products obtainable by curing such a composition and to the use of such a composition as electrical insulating material.

Epoxy resins are frequently used as electrical insulating material on account of their good mechanical properties and their high specific resistance. Cycloaliphatic epoxy resins are especially suitable for outdoor applications because of their high resistance to weathering, but the problem arises, especially in regions having high levels of precipitation and air pollution, that a conductive dirt/water layer can form on the surface of the insulator, which leads to leakage currents and arcing and may have consequences ranging from damage to the insulator up to total failure. Even only slightly soiled epoxy-based insulators can undergo a rise in surface conductivity if the surface becomes eroded by weathering in the course of time and the water is better able to wet the resulting roughened layer.

As disclosed in US Patent 3 926 885, epoxy resins can be provided with hydrophobic properties by the addition of polysiloxane/polyether copolymers and OH-terminated polysiloxanes, but the adhesion of the resulting material to metal is not sufficient for all applications.

JP-A 2-305454 describes epoxy resin mixtures having a high degree of stability towards moisture which, in addition to comprising an epoxy novolak and a phenolic resin, also comprise small amounts of a cyclic dimethylsiloxane. Although, in those compositions, the corrosion caused by binding of water on the surface is largely prevented, a hydrophobicity effect sufficient for use as an insulator is not achieved with such systems.

WO 98/32138 describes a resin system suitable as electrical insulating material that is based on curable mixtures of epoxy resins and specific silicone oligomers having terminal glycidyl groups. As a result of the curing, the silicone oligomers become part of the crosslinked structure that is formed, so that known properties of silicones, such as hydrophobicity and good resistance to weathering, can be imparted to the cured material. A disadvantage is the use of expensive commercially available silicone oligomers and a poor hydrophobicity transfer effect.

It has now been found that compositions comprising an epoxy resin, at least two specific polysiloxanes and a non-ionic, fluoroaliphatic surface-active reagent are able to yield storage-stable emulsions that in the fully cured state have a pronounced hydrophobicity transfer effect and recovery effect.

The present invention relates to a composition comprising

- (a) an epoxy resin,
- (b) an OH-terminated polysiloxane,
- (c) a cyclic polysiloxane and
- (d) a non-ionic, fluoroaliphatic surface-active reagent.

The amounts of components (a) to (c) in the compositions according to the invention can vary within wide limits.

Preference is given to compositions comprising,

based on the total composition (a), (b), (c) and (d),

from 77.0 to 97.99 % by weight, especially from 86.0 to 96.95 % by weight, component (a), from 1.0 to 10.0 % by weight, especially from 2.0 to 6.0 % by weight, component (b), from 1.0 to 10.0 % by weight, especially from 1.0 to 5.0 % by weight, component (c) and from 0.01 to 3.0 % by weight, especially from 0.05 to 3.0 % by weight, component (d), the sum of components (a), (b), (c) and (d) being 100 % by weight.

As component (a) of the compositions according to the invention there are suitable any type of epoxide that contains at least one glycidyl or β-methylglycidyl group, a linear alkylene oxide group or a cycloalkylene oxide group.

Examples of suitable epoxy resins are polyglycidyl and poly(β -methylglycidyl) ethers obtainable by reaction of a compound containing at least two free alcoholic and/or phenolic hydroxyl groups per molecule with epichlorohydrin or β -methylepichlorohydrin under alkaline conditions, or alternatively in the presence of an acid catalyst with subsequent alkali treatment.

Suitable starting compounds for the preparation of such glycidyl or β-methylglycidyl ethers are, for example, acyclic alcohols, such as ethylene glycol, diethylene glycol and higher poly(oxyethylene) glycols, propane-1,2-diol and poly(oxypropylene) glycols, propane-1,3-diol,

butane-1,4-diol, poly(oxytetramethylene) glycols, pentane-1,5-diol, hexane-1,6-diol, hexane-2,4,6-triol, glycerol, 1,1,1-trimethylolpropane, pentaerythritol or sorbitol, cycloaliphatic alcohols, such as resorcitol, quinitol, bis(4-hydroxycyclohexyl)methane, 2,2-bis(4-hydroxycyclohexyl)propane and 1,1-bis(hydroxymethyl)cyclohex-3-ene, and alcohols having aromatic nuclei, such as N,N-bis(2-hydroxyethyl)aniline and p,p'-bis(2-hydroxyethylamino)diphenyl-methane.

Further suitable dihydroxy compounds for the preparation of glycidyl or β -methylglycidyl ethers are mononuclear phenols, such as resorcinol and hydroquinone, polynuclear phenols, such as bis(4-hydroxyphenyl)methane, 4,4-dihydroxydiphenyl, bis(4-hydroxyphenyl)sulfone, 1,1,2,2-tetrakis(4-hydroxyphenyl)ethane, 2,2-bis(4-hydroxyphenyl)propane (bisphenol A) and 2,2-bis(3,5-dibromo-4-hydroxyphenyl)propane, and novolaks, for example phenol and cresol novolaks.

Polyglycidyl and poly(β-methylglycidyl) esters are obtainable by reaction of a compound containing two or more carboxylic acid groups per molecule with epichlorohydrin, glycerol dichlorohydrin or β-methylepichlorohydrin in the presence of alkali. Such polyglycidyl esters can be derived from aliphatic polycarboxylic acids, e.g. oxalic acid, succinic acid, glutaric acid, adipic acid, pimelic acid, suberic acid, azelaic acid, sebacic acid or dimerised or trimerised linoleic acid, from cycloaliphatic polycarboxylic acids, such as tetrahydrophthalic acid, 4-methyltetrahydrophthalic acid, hexahydrophthalic acid and 4-methylhexahydrophthalic acid, and from aromatic polycarboxylic acids, such as phthalic acid, isophthalic acid and terephthalic acid.

Further epoxides suitable as component (a) are poly(N-glycidyl) compounds, for example the products obtainable by dehydrochlorination of the reaction products of epichlorohydrin and amines containing at least two amino hydrogen atoms, such as aniline, n-butylamine, bis(4-aminophenyl)methane and bis(4-methyl-aminophenyl)methane. Also included are triglycidyl isocyanurate as well as N,N'-diglycidyl derivatives of cyclic alkyleneureas, such as ethyleneurea and 1,3-propyleneurea and hydantoins, such as 5,5-dimethylhydantoin. Poly(S-glycidyl) compounds, for example the di-S-glycidyl derivatives of dithiols, such as ethane-1,2-dithiol and bis(4-mercaptomethylphenyl) ether, are likewise suitable.

Preferably the compositions comprise as component (a) a cycloaliphatic epoxy resin or an epoxidation product of a natural unsaturated oil or a derivative thereof.

The term "cycloaliphatic epoxy resin" in the context of this invention denotes any epoxy resin having cycloaliphatic structural units, that is to say it includes both cycloaliphatic glycidyl compounds and β -methylglycidyl compounds as well as epoxy resins based on cycloalkylene oxides.

Suitable cycloaliphatic glycidyl compounds and β -methylglycidyl compounds are the glycidyl esters and β -methylglycidyl esters of cycloaliphatic polycarboxylic acids, such as tetrahydrophthalic acid, 4-methyltetrahydrophthalic acid, hexahydrophthalic acid, 3-methylhexahydrophthalic acid and 4-methylhexahydrophthalic acid.

Further suitable cycloaliphatic epoxy resins are the diglycidyl ethers and β–methylglycidyl ethers of cycloaliphatic alcohols, such as 1,2-dihydroxycyclohexane, 1,3-dihydroxycyclohexane and 1,4-dihydroxycyclohexane, 1,4-cyclohexanedimethanol, 1,1-bis(hydroxymethyl)-cyclohex-3-ene, bis(4-hydroxycyclohexyl)methane, 2,2-bis(4-hydroxycyclohexyl)propane and bis(4-hydroxycyclohexyl)sulfone.

Examples of epoxy resins having cycloalkylene oxide structures are bis(2,3-epoxycyclopentyl) ether, 2,3-epoxycyclopentylglycidyl ether, 1,2-bis(2,3-epoxycyclopentyl)ethane, vinyl cyclohexene dioxide, 3,4-epoxycyclohexylmethyl 3',4'-epoxycyclohexanecarboxylate, 3,4-epoxy-6-methylcyclohexylmethyl 3',4'-epoxy-6'-methylcyclohexanecarboxylate, bis(3,4-epoxycyclohexylmethyl) adipate and bis(3,4-epoxy-6-methylcyclohexylmethyl) adipate.

Preferred cycloaliphatic epoxy resins are bis(4-hydroxycyclohexyl)methanediglycidyl ether, 2,2-bis(4-hydroxycyclohexyl)propanediglycidyl ether, tetrahydrophthalic acid diglycidyl ester, 4-methyltetrahydrophthalic acid diglycidyl ester, 4-methylhexahydrophthalic acid diglycidyl ester, and especially hexahydrophthalic acid diglycidyl ester and 3,4-epoxycyclohexylmethyl 3',4'-epoxycyclohexanecarboxylate.

As component (a) it is also possible to use in the compositions according to the invention epoxidation products of unsaturated fatty acid esters. It is preferable to use epoxy-containing compounds derived from mono- and poly-fatty acids having from 12 to 22 carbon atoms and an iodine number of from 30 to 400, for example lauroleic acid, myristoleic acid.

palmitoleic acid, oleic acid, gadoleic acid, erucic acid, ricinoleic acid, linoleic acid, linolenic acid, elaidic acid, licanic acid, arachidonic acid and clupanodonic acid.

For example, there are suitable the epoxidation products of soybean oil, linseed oil, perilla oil, tung oil, oiticica oil, safflower oil, poppyseed oil, hemp oil, cottonseed oil, sunflower oil, rapeseed oil, polyunsaturated triglycerides, triglycerides from euphorbia plants, groundnut oil, olive oil, olive kernel oil, almond oil, kapok oil, hazelnut oil, apricot kernel oil, beechnut oil, lupin oil, maize oil, sesame oil, grapeseed oil, lallemantia oil, castor oil, herring oil, sardine oil, menhaden oil, whale oil, tall oil and derivatives thereof.

Also suitable are higher unsaturated derivatives that can be obtained by subsequent dehydrogenation reactions of those oils.

The olefinic double bonds of the unsaturated fatty acid radicals of the above-mentioned compounds can be epoxidised in accordance with known methods, for example by reaction with hydrogen peroxide, optionally in the presence of a catalyst, an alkyl hydroperoxide or a per acid, for example performic acid or peracetic acid.

Within the scope of the invention, both the fully epoxidised oils and the partially epoxidised derivatives that still contain free double bonds can be used as component (a).

Especially preferred as component (a) are epoxidised soybean oil and epoxidised linseed oil.

The OH-terminated polysiloxanes in accordance with component (b) can be prepared according to known methods, for example by hydrolysis of the corresponding organochlorosilanes and subsequent polycondensation of the silanols, there generally being formed polysiloxane mixtures having molecular masses of from 1000 to 150 000 g/mol. A number of such OH-terminated polysiloxanes are commercially available.

In the compositions according to the invention it is preferable to use liquid polysiloxanes.

There is preferably used a polysiloxane of formula I

$$HO = \begin{bmatrix} R_1 \\ Si - O \end{bmatrix} H \qquad (I),$$

$$R_2 = \begin{bmatrix} R_1 \\ R_2 \end{bmatrix} R$$

wherein R_1 and R_2 are each independently of the other C_1 - C_{18} alkyl, C_5 - C_{14} aryl or C_6 - C_{24} -aralkyl and n is an average value of from 3 to 60, especially from 4 to 20.

Alkyl includes, for example, methyl, ethyl, isopropyl, n-propyl, n-butyl, isobutyl, sec-butyl, tert-butyl and the various isomeric pentyl, hexyl, heptyl, octyl, nonyl, decyl, undecyl, dodecyl, tridecyl, tetradecyl, pentadecyl, hexadecyl, heptadecyl, octadecyl, nonadecyl and eicosyl groups.

Aryl as R_1 or R_2 contains preferably from 6 to 14 carbon atoms and may be, for example, phenyl, tolyl, pentalinyl, indenyl, naphthyl, azulinyl and anthryl.

Aralkyl as R_1 or R_2 contains preferably from 7 to 12 carbon atoms and especially from 7 to 10 carbon atoms. It may be, for example, benzyl, phenylethyl, 3-phenylpropyl, α -methylbenzyl, 4-phenylbutyl or α , α -dimethylbenzyl.

Special preference is given to polysiloxanes of formula I wherein R_1 and R_2 are each independently of the other methyl, ethyl or phenyl.

Especially preferred as component (b) are polysiloxanes of formula I wherein R_1 and R_2 are methyl and n = 4 to 20.

The cyclic polysiloxanes in accordance with component (c) are likewise known to the person skilled in the art and can be prepared according to known methods.

There is preferably used as component (c) a cyclic polysiloxane of formula II

$$\begin{array}{c|c}
R_1 \\
Si - O \\
R_2
\end{array}$$
(II),

wherein R_1 and R_2 are each independently of the other C_1 - C_{18} alkyl, C_5 - C_{14} aryl or C_6 - C_{24} -aralkyl and m is an integer from 3 to 12.

Alkyl, aryl and aralkyl in formula (II) have the same definitions as the corresponding groups according to formula (I).

Preferred as component (c) are cyclic polysiloxanes of formula II wherein R_1 and R_2 are each independently of the other methyl, ethyl or phenyl and m is an integer from 3 to 8.

Very especially, R₁ and R₂ are methyl and m is an integer from 6 to 8.

As described in J. Am. Chem. Soc. <u>68</u>, 358 (1946), such cyclic polysiloxanes can be isolated from the product mixture formed on hydrolysis of the corresponding dialkyl-, diaryl- or diaralkyl-dichlorosilanes.

Especially preferred as component (c) are the commercially available compounds octamethylcyclotetrasiloxane (m = 4), decamethylcyclopentasiloxane (m = 5) and especially dodecamethylcyclohexasiloxane (m = 6) and also hydrolysates of dimethyldichlorosilane, especially undistilled hydrolysates, since they have higher contents of cyclic polysiloxanes having the preferred ring size m = 6 to 8, and hence in addition to dodecamethylcyclohexasiloxane also tetradecamethylcycloheptasiloxane (m = 7) and hexadecamethylcyclooctasiloxane (m = 8).

As surface-active reagent in the form of component (d) of the compositions according to the invention there are suitable derivatives of non-ionic, perfluorinated polyalkylenes, such as perfluorinated polyoxyalkylenes. Preference is given to compounds of a combination of a perfluorinated aliphatic alkyl unit R_f with a hydrocarbon unit R_f , in which the latter contains at least one, preferably oxygen-containing, mono- or di-valent polar functional group, e.g. -OH, -COOH, -COOR, -COO-, -CO-, -O-.

Suitable compounds are alkoxylated, especially ethoxylated, perfluorinated fatty acid derivatives, for example:

$$R_f$$
 -COO-(CH_2CH_2O)_m-R (III)

or

$$R_f - (CH_2CH_2O)_m - R$$
 (IV),

wherein m = 1 to 200, R_f is a linear or branched perfluorinated alkyl having from 2 to 22 carbon atoms and R = H, C_1 - C_6 alkyl or R_f .

It is preferable to use compounds of formula (III) or (IV) in which the molecular mass according to the theoretical empirical formula is only from 200 to 10 000, especially from 300 to 8000.

Preferred compounds are, for example, F_3C -(CF_2)₅-(CH_2CH_2O)-H = 1,1,2,2-tetrahydro-perfluorooctanol (corresponding to formula (IV), wherein R_f = perfluorinated n-hexyl, m = 1 and R = H) or R_f -COO-(CH_2CH_2O)_m-R, wherein R_f is a linear perfluorinated alkyl having from 16 to 18 carbon atoms, m = 110-130 and R = H.

Some of those compounds are commercially obtainable from a number of sources, e.g. as ZONYL® Fluorochemical Intermediates (DuPont), e.g. ZONYL® BA-L and BA Fluoroalcohols, or FLUORAD® Fluorosurfactants (3M), e.g. FLUORAD® FC-431. Further surfaceactive compounds suitable for use according to the invention can be found in the technical bulletins of the manufacturers mentioned, for example "Technical Information" 233592B (1/94) relating to ZONYL® Fluorochemical Intermediates (DuPont).

One or more compounds of each component (a) to (d) can be used in the compositions according to the invention.

The compositions according to the invention can be cured in principle by cationic polymerisation of the epoxy resin system using an initiator system or using any customary epoxy hardener, but it is preferable to use anhydride hardeners.

The invention therefore relates also to a composition comprising the above-mentioned components (a) to (d) and additionally as component (e) a polycarboxylic anhydride.

Such an anhydride may be a linear aliphatic polymeric anhydride, for example polysebacic polyanhydride or polyazelaic polyanhydride, or a cyclic carboxylic anhydride.

Cyclic carboxylic anhydrides are especially preferred.

Examples of cyclic carboxylic anhydrides are:

succinic anhydride, citraconic anhydride, itaconic anhydride, alkenyl-substituted succinic anhydrides, dodecenylsuccinic anhydride, maleic anhydride and tricarballylic anhydride, a

maleic anhydride adduct with cyclopentadiene or methylcyclopentadiene, a linoleic acid adduct with maleic anhydride, alkylated endoalkylenetetrahydrophthalic anhydrides, methyltetrahydrophthalic anhydride and tetrahydrophthalic anhydride, the isomeric mixtures of the two latter compounds being especially suitable. Especially preferred are hexahydrophthalic anhydride and methylhexahydrophthalic anhydride.

Further examples of cyclic carboxylic anhydrides are aromatic anhydrides, for example pyromellitic dianhydride, trimellitic anhydride and phthalic anhydride.

It is also possible to use chlorinated or brominated anhydrides, e.g. tetrachlorophthalic anhydride, tetrabromophthalic anhydride, dichloromaleic anhydride and chlorendic anhydride.

The compositions according to the invention can optionally additionally comprise a curing accelerator (f). Suitable accelerators are known to the person skilled in the art. Examples that may be mentioned are:

complexes of amines, especially tertiary amines, with boron trichloride or boron trifluoride; tertiary amines, such as benzyldimethylamine;

urea derivatives, such as N-4-chlorophenyl-N',N'-dimethylurea (monuron); unsubstituted or substituted imidazoles, such as imidazole or 2-phenylimidazole.

Preferred accelerators are tertiary amines, especially benzyldimethylamine, and imidazoles (e.g. 1-methylimidazole) for the above-mentioned compositions that comprise epoxidised oils.

The components (e) and (f) are used in the customary effective amounts, that is to say amounts sufficient for curing the compositions according to the invention. The ratio of components (a) and (e) and optionally (f) depends upon the nature of the compounds used, the rate of curing required and the properties desired in the end product and can readily be determined by the person skilled in the art. Generally, from 0.4 to 1.6 equivalents, preferably from 0.8 to 1.2 equivalents, of anhydride groups per epoxy equivalent are used.

The resin mixture (a) to (d) and the hardener component (e), optionally together with the accelerator (f), are generally stored separately and mixed together only shortly before

application. If the resin mixture (a) to (d) is to be kept in interim storage prior to curing, the resin mixture (a) to (d) requires an additional auxiliary as an optional component (g) in order that the mixture, which forms an emulsion, can be maintained in storable form. As such a stabilising auxiliary there may be used emulsifiers (surface-active and interfacially active compounds) and thickeners (for example silicic acids, bentonites, dibenzylidenesorbitol etc.). Such auxiliaries and their use are well known to the person skilled in the art.

Of the auxiliaries mentioned, the use of highly dispersed silicic acid is preferred. Highly dispersed, hydrophilic, untreated silicic acids are especially suitable. They are commercially available, for example in the form of Aerosil[®]. The effective amounts of silicic acid are in the range of from 0.01 to 3.5 % by weight, preferably from 0.05 to 3.0 % by weight, based on the sum of components (a) to (d), and the average size of the primary particles is advantageously about 12 nm.

The invention therefore relates also to storage-stable compositions obtainable by the addition of stabilising auxilaries, such as emulsifiers and thickeners. Instead of hardener component (e), optionally together with an accelerator (f), the resin mixture (a) to (d) may comprise as component (e) an initiator system for the cationic polymerisation of the epoxy resin. As initiator system for the cationic polymerisation of the epoxy resins there are used, for example, thermally activatable initiators, such as thermally activatable onium salts, oxonium salts, iodonium salts, sulfonium salts, phosphonium salts or quaternary ammonium salts that do not contain nucleophilic anions. Such initiators and their use are known. For example, US Patent 4 336 363, EP-A-0 379 464 and EP-A-0 580 552 disclose specific sulfonium salts as curing agents for epoxy resins. US Patent 4 058 401, in addition to describing specific sulfonium salts, also describes the corresponding salts of tellurium and selenium.

Quaternary ammonium salts as thermally activatable initiators are disclosed, for example, in EP-A-0 066 543 and in EP-A-0 673 104. They are salts of aromatic-heterocyclic nitrogen bases with non-nucleophilic, for example complex, halide anions, such as BF_4^- , PF_6^- , SbF_6^- , $SbF_5(OH)^-$ and AsF_6^- .

As quaternary ammonium salt there is especially used N-benzylquinolinium hexafluoroantimonate. When quaternary ammonium salts are used it is advantageous to use in addition a thermal free-radical former, for example pinacols and their ethers, esters or silyl derivatives. Such compounds are known and can be prepared in accordance with known procedures.

As thermal free-radical formers there are preferably used the pinacols, such as acetophenone pinacols or especially 1,1,2,2-tetraphenyl-1,2-ethanediol (benzopinacol).

As thermally activatable initiator there is especially used N-benzylquinolinium hexafluoroantimonate together with 1,1,2,2-tetraphenyl-1,2-ethanediol, preferably in a molar ratio of about 1:1.

The activation temperature of the cationic initiators is generally above room temperature, preferably in the range of from 60 to 180°C, especially from 90 to 150°C.

The amount of cationic initiator contained in the cationically curable epoxy resin is generally from 0.05 to 30 % by weight, preferably from 0.5 to 15 % by weight, based on the amount of cationically polymerisable epoxy resin.

Furthermore, the curable mixtures may comprise tougheners, for example core/shell polymers or the elastomers or elastomer-containing graft polymers known to the person skilled in the art as rubber tougheners.

Suitable tougheners are described, for example, in EP-A-0 449 776. They are preferably used in an amount of from 1 to 20 % by weight, based on the total amount of epoxy resin in the composition.

The curable mixtures may also comprise further fillers in addition to those mentioned above, for example metal powder, wood flour, glass powder, glass beads, semi-metal and metal oxides, such as SiO₂ (quartz sand, quartz powder, silanised quartz powder, fused silica powder, silanised fused silica powder), aluminium oxide, titanium oxide and zirconium oxide, metal hydroxides, such as Mg(OH)₂, Al(OH)₃, silanised Al(OH)₃ and AlO(OH), semi-metal and metal nitrides, for example silicon nitride, boron nitrides and aluminium nitride, semi-metal and metal carbides (SiC and boron carbides), metal carbonates (dolomite, chalk, CaCO₃), metal sulfates (barite, gypsum), ground minerals, e.g. of hydromagnesite and huntite, and natural or synthetic minerals chiefly of the silicate series, e.g. zeolites (especially

molecular sieves), talcum, mica, kaolin, wollastonite and others. Preferred fillers are quartz powder, silanised quartz powder, aluminium hydroxide and aluminium oxide.

In addition to the additives mentioned above, the curable mixtures may also comprise further customary ingredients, e.g. antioxidants, light stabilisers, flame retardants, fillers containing water of crystallisation, plasticisers, dyes, pigments, fungicides, thixotropic agents, toughness improvers, antifoams, antistatics, lubricants, anti-settling agents, wetting agents and mould-release agents.

The compositions according to the invention can be produced in accordance with known methods using known mixing apparatus, for example stirrers (especially dispersers and Supraton® having a high shear gradient), kneaders, rollers or dry mixers. In the case of solid epoxy resins, the dispersing can also be carried out in the melt.

The curing of the mixtures according to the invention can be carried out in known manner in one or more stages. It is generally effected by heating the mixtures to temperatures of from 60°C to 200°C, especially from 80°C to 180°C.

The invention relates also to the crosslinked products obtainable by curing a composition according to the invention.

Surprisingly, in comparison with the corresponding unmodified systems (Comparison Example 1), the addition of the two chemically different siloxane components and the surface-active reagent in the compositions according to the invention results in virtually no impairment, or only slight impairment, of the mechanical and electrical properties of the products obtained therefrom.

The addition of silicones generally results in impairment of the adhesion properties, but the compositions according to the invention nevertheless unexpectedly exhibit good adhesion to metal, which is revealed by the fact that the good cantilever strength and pull-out strength values remain unchanged. Likewise, shaped articles encapsulated using a system according to the invention are surprisingly found to have a comparatively very good temperature change behaviour.

In comparison with unmodified systems, the invention offers the advantage that traces of the siloxanes used according to the invention are able to migrate onto any soiling present on the

fully cured material. As a result, an initially hydrophilic layer of dirt also becomes hydrophobic (hydrophobicity transfer), with the result that water more readily rolls away from the soiling in the form of beads and does not form a coherent conductive, and therefore harmful, dirt/water layer, as is the case with the unmodified system. This effect is even, surprisingly, very pronounced. The water runs off in a considerably better and more rapid manner. Surprisingly, the durability of this effect is also very good, that is to say the pronounced hydrophobicity transfer effect is retained even when the layer of dirt is repeatedly removed and reapplied.

Even in the case of only slightly soiled epoxy-resin-based insulators, there may be a loss of the original hydrophobicity and thus a rise in the surface conductivity, the reason being microdischarges which can be caused, for example, by individual raindrops on the insulator surface.

In comparison with unmodified systems (Comparison Example 1), a further advantage of the present invention is that, by means of the present systems, the hydrophobicity may surprisingly be regained after such an imposed loss, that is to say the original hydrophobicity is restored within a period ranging from hours to a few days ("recovery" effect). As a result, the systems according to the invention with their excellent hydrophobicity properties in the form of a very good and at the same time long-lasting hydrophobicity transfer effect in combination with a very good recovery effect and good temperature change properties are by their nature suitable for use as insulating material for outdoor applications in fields of use where climatic conditions are difficult.

The compositions according to the invention are especially suitable as casting resins, casting compounds ("structural casting"), laminating resins, compression moulding compounds ("epoxy moulding compounds"), coating compounds and especially as electrical insulating compounds.

The invention relates also to the use of the compositions according to the invention as electrical insulating material.

In the following Examples, the following commercially available substances are used:

epoxy resin 1: liquid hexahydrophthalic acid diglycidyl ester; epoxy content: 5.6 to 6.2 eq./kg ("CY 184", Ciba Spezialitäten Chemie) ESO epoxidised soybean oil; epoxy content: 4.10 to 4.20 eq./kg;

("Reoplast", Witco)

ELO epoxidised linseed oil; epoxy content: 5.50 to 5.65 eq./kg:

("Merginat", Harburger Chemie)

hardener 1 hardener mixture of 70 parts by weight of hexahydrophthalic anhydride and

30 parts by weight of methylhexahydrophthalic anhydride

W 12: untreated quartz powder (Quarzwerke Frechen)

W 12 EST: quartz powder pretreated with epoxysilane (Quarzwerke Frechen)

polysiloxane 1: OH-terminated polydimethylsiloxane having a viscosity of 5 Pa · s

("NG200-5000", Wacker)

polysiloxane 2: mixture of linear OH-terminated (≤ 40%) polydimethylsiloxanes and cyclic

dimethylsiloxanes (≥ 60%) having a viscosity of 5 to 20 Pa · s ("Dimethyl-

methanolysat", GE-Bayer AG)

Fluorad: non-ionic perfluorinated aliphatic polymeric ester (ethoxylated

perfluorinated fatty acid) ("Fluorad FC 431", 3M)

BDMA benzyldimethylamine

1-MI 1-methylimidazole

Aerosil hydrophilic, highly dispersed silicic acid ("Aerosil 200", Degussa)

Preparation Examples

Comparison Example 1:

In a mixing apparatus having a blade stirring mechanism, 1000 g of epoxy resin 1, 900 g of hardener 1, 5.0 g of BDMA and 2700 g of quartz powder W 12 EST are mixed together within a period of 30 min at 60°C using a mixer having a blade stirring mechanism and then briefly degassed at about 10 mbar. The composition is then cured for 6 hours at 80°C and for 10 hours at 140°C. The properties of the cured product are summarised in Table 1.

Invention Example 1:

In a mixing apparatus having a disperser disc, 9050 g of epoxy resin 1, 200 g of polysiloxane 1, 500 g of polysiloxane 2 and 100 g of Fluorad are mixed together at 3750 rev/min at room temperature in the course of 10 minutes. 150 g of Aerosil are then added to the mixture and mixing is carried out at 3750 rev/min at room temperature within a period of one hour, yielding a white, storage-stable emulsion.

1000 g of the resin premix so prepared are mixed with 814.5 g of hardener 1, 4.5 g of BDMA and 2578.1 g of quartz powder W 12 EST within a period of 30 min at 60°C using a mixer having a blade stirring mechanism and then briefly degassed at about 10 mbar. The composition is then cured for 6 hours at 80°C and for 10 hours at 140°C. The properties of the cured product are summarised in Table 1.

<u>Invention Example 2:</u>

In a mixing apparatus having a disperser disc, 6855 g of epoxy resin 1, 200 g of polysiloxane 1, 500 g of polysiloxane 2, 10 g of Fluorad, 1143 of ESO and 1143 g of ELO are mixed together at 3750 rev/min at room temperature in the course of 10 minutes. 150 g of Aerosil are then added to the mixture and mixing is carried out at 3750 rev/min at room temperature within a period of one hour, yielding a white, storage-stable emulsion. 1000 g of the resin premix so prepared are mixed with 806.6 g of hardener 1, 2.3 g of BDMA, 4.5 g of 1-MI and 2567.1 g of quartz powder W 12 EST within a period of 30 min at 60°C using a mixer having a blade stirring mechanism and then briefly degassed at about 10 mbar.

The composition is then cured for 6 hours at 80°C and for 10 hours at 140°C. The properties of the cured product are summarised in Table 1.

Invention Example 3:

In a mixing apparatus having a disperser disc, first of all 50 g of dibenzylidenesorbitol are dissolved in 9100 g of epoxy resin 1 at a maximum of 110° and the mixture is then cooled to room temperature. 200 g of polysiloxane 1, 500 g of polysiloxane 2 and 100 g of Fluorad are then added and the mixture is stirred at 1500 rev/min at room temperature for 15 minutes. 50 g of Aerosil are then added to the mixture and at room temperature the entire mixture is stirred at 1500 rev/min for a further one hour, yielding a white, storage-stable emulsion. 1000 g of the resin premix so prepared are mixed with 820 g of hardener 1, 4.5 g of BDMA and 3542 g of quartz powder W 12 EST within a period of 30 min at 60°C using a mixer having a blade stirring mechanism and then briefly degassed at about 10 mbar.

Application Example 1:

Pin insulators are produced by means of the pressure gelation process. For that purpose, the uncured mixtures, prepared according to Example 1 and analogously to Comparison

Example 1 but with a degree of filler of 66 % by weight, are injected into a metal mould which has been heated to 140°C and treated with parting agent. After gelation (after about 20 min), the casting is removed from the mould and post-cured for 10 hours at 140°C. The insulator produced from the composition according to the invention surprisingly has a cantilever strength of a level similar to that of the insulator produced from the analogous composition without silicone additives, while a comparison of the pull-out strengths shows that the material according to the invention still adheres very well to the insert (see Table 1).

Application Example 2:

The improved behaviour of the insulators produced using the system modified according to the invention in atmospheres having a high level of air pollution, especially the improved hydrophobicity properties, are demonstrated by the following tests:

2A: Hydrophobicity transfer effect

Specimens of Invention Examples $\underline{1}$ to $\underline{3}$ are tested to demonstrate the so-called "hydrophobicity transfer effect". For that purpose, artificial soiling is applied to \underline{four} plates made of material prepared with the product according to Invention Examples $\underline{1}$ to $\underline{3}$ and Comparison Example 1 (prepared without the use of silicone-containing parting agents and cleaned with acetone after demoulding). For that purpose, quartz powder W 12 is applied to the plates from a vibrating apparatus in an amount per unit area sufficient to produce a foreign layer 0.5 mm thick when the powder is smoothed with a ruler or the like. In order to test whether or not the material then transfers the hydrophobicity to the quartz layer, which is actually hydrophilic, at specific time intervals a 30 μ l water droplet is applied to the foreign layer using a pipette, the behaviour of the droplet is observed and classified in accordance with the following scheme:

Transfer state system (TS)

TS	Properties	Level specimen	Specimen inclined at 45°	Specimen initially level, then inclined at 45°
1	No hydro- phobicity transfer effect (HTE)	Droplet is immediately absorbed		
1.5		Droplet is absorbed within 2 minutes		
2	weak HTE	Droplet remains stable for 2 minutes		
2.5			Droplet forms "nose shape"	
3	moderate HTE		Runs off with a large amount of foreign layer material	Droplet remains stable on tilting
3.5			Runs off with a large amount of foreign layer material	Droplet forms "nose shape" on tilting
4	good HTE			Droplet runs off on tilting, leaving a clearly visible track
4.5				Droplet leaves a slight track on tilting
5	excellent HTE	Droplet runs off without foreign material and "dances about" when specimen is tilted slightly		

Transfer state system: $500 \, \mu m$ thick quartz powder layer (quartz powder W12, Quarzwerke Frechen), Application $30 \, \mu l$ water droplet

2B: Demonstration of the "recovery" effect by means of a plasma test

In order to demonstrate the recovery effect, the state of hydrophobicity is determined (see 2B1). Then a state of hydrophilicity is imposed by the use of a plasma (see 2B2). After the plasma treatment, which is intended to bring about loss of hydrophobicity, the state of the surface is again investigated at various time intervals (immediately, after one hour (h) and one day (d)). A recovery effect is present when a specimen passes from a hydrophilic state (in about class 5 to 7) to a hydrophobic state (in about class 1 to 4), that is to say when the state prior to the plasma treatment has approximately been restored. The results are summarised in Table 1.

<u>2B1</u>: The hydrophobicity state of the specimens at any given time is determined as follows: a vertically arranged specimen surface about 100 cm^2 in size is sprayed with water 20 times (once per second) from a distance of about $25 \pm 10 \text{ cm}$ using a spray bottle. After a further

10 seconds the specimen is examined and the state of the surface is classified in accordance with the following scheme:

Classification of the hydrophobicity state in the spraying test

Class HC	Properties	Description of the effects
1	Excellent surface hydrophobicity	Only discrete droplets, contact angle for the majority of droplets is > 80°
2	Very good surface hydrophobicity	Only discrete droplets, contact angle for the majority of droplets is > 50° but < 80°
3	Good surface hydrophobicity	Only discrete droplets, contact angle for the majority of droplets is > 20° but < 50°
4	Moderate surface hydrophobicity	Discrete droplets and discrete wetted areas, the size of the individual wetted areas is < 2 cm ² and the total wetted area is < 90%
5	Low surface hydrophobicity	Some discrete wetted areas, areas > 2 cm ² , the size of the total wetted area is < 90%
6	Low hydrophilicity	> 90% of the total area is wetted but some unwetted sites can be identified
7	Very hydrophilic surface	Continuous film of water over the entire area tested

<u>2B2</u>: A loss of hydrophobicity, i.e. hydrophilicity, is imposed by means of a plasma treatment. For that purpose, the test plates, which have dimensions of 10 x 10 x 0.4 cm³, are placed in a plasma chamber type 440 (Technics Plasma GmbH) and exposed to the following conditions: plasma treatment time: 2 min, pressure: 2 to 3 mbar, gas: oxygen, output: 200 Watt. The treatment results in the loss of hydrophobicity (see Table 1).

Comparative Application Examples:

Three Comparison Examples are prepared in principle analogously to Invention Example 1. Unlike Invention Example 1, in which polysiloxanes 1 and 2 and a surface-active reagent are used at the same time, in each Comparison Example only one additive is used. The individual compositions and the properties of the cured products are summarised in Table 1.

Table 1

CE 1	IE 1	IE 2	IE3	CE 2	CE 3	CE 4
100	90.5	68.55	91	97	94	99
ļ		11.43				
		11.43				
	2	2	2	2		
	5	5	5	}	5	
	1	0.1	1			1
0.5	0.45	0.23	0.45	0.49	0.47	0.5
		0.45			ļ	
	1.5	1.5	0.5	1	1	
			0.5	ļ		
90	81.45	80.66	82	87.3	84.6	89.1
270	257.8	254.6	354.2	266.1	262.3	268.7
one	good	good	good	poor	poor	one
phase						phase
110	110	109	110			
1	0.5	0.2				
2	1.6	1.7				
125	127	123				
90	86	75				
	1.4					
	137					
	1.6	1.7				
10500	9785	8813				
	2.24	2.12				
485	465	465				
۸۶۵	45.0	45.0	A.F.O.	450	45.0	45.0
					AE 2	AE 2
						1
				3.5		1
			4.5			
						2
						6
						7
ь	1	ა		5	5	5
AE 1	AE 1	AE1	AE3			
66	66	66	66			
6125	5972	5509	crack			
71.6	64.8	62	test			
	100 0.5 90 270 one phase 110 1 2 125 90 1.4 125 1.3 10500 2.25 485 AE 2 1 1 3 7 7 6 AE 1 66 6125	100 90.5 2 5 1 0.5 0.45 1.5 90 81.45 270 257.8 one phase good phase 110 110 1 0.5 2 1.6 125 127 90 86 1.4 1.4 125 137 1.3 1.6 10500 9785 2.25 2.24 485 465 AE 2 AE 2 1 3.5 1 4 1 4 3 2 7 7 7 2 6 1 AE 1 AE 1 66 6125 5972	100 90.5 68.55 11.43 11.43 11.43 11.43 11.43 11.43 11.43 11.43 11.43 11.43 11.43 0.1 0.45 0.23 0.45 1.5 90 81.45 80.66 270 257.8 254.6 one phase good good 110 110 109 1 0.5 0.2 2 1.6 1.7 125 127 123 90 86 75 1.4 1.4 1.3 125 137 127 1.3 1.6 1.7 10500 9785 8813 2.25 2.24 2.12 485 465 465 AE 2 AE 2 AE 2 1 4 4 4 4 4 4 4 4 4 4 4 4	100 90.5 68.55 11.43 11.43 11.43 11.43 11.43 11.43 11.43 11.43 11.43 11.43 11.43 11.43 11.43 11.43 11.5 1.5 1.5 1.5 1.5 1.5 1.5 1.5 1.5 1.	100 90.5 68.55 91 97 11.43 11.43 2 2 2 2 5 5 5 5 5 1 0.1 1 1 0.5 0.45 0.45 0.45 0.45 0.45 0.5 1 0.5	100 90.5 68.55 91 97 94 1.1.43 11.43 2 2 2 5 5 5 5 5 5 5 5 5 5 5 5 5 5 5 5

⁻ DSC (Differential Scanning Calorimetry) carried out using a TA 4000 apparatus (Mettler)

⁻ electrical values (tangent δ) according to DIN 53483, measurement frequency 50 Hz

⁻ tensile strength and breaking strength according to ISO R527

⁻ bending strength, outer fibre strain and elasticity modulus in flexure according to ISO 178

⁻ K1C and G1C: double torsion test

Table 1 shows:

1) The properties of an unmodified reference material:

No hydrophobicity transfer is found, but good surface hydrophobicity. After the loss thereof (by means of plasma), however, the surface hydrophobicity is not restored.

2) The properties of the modified compositions according to the invention:

Example 1 very quickly exhibits a good hydrophobicity transfer effect, as it does also in the second application of the foreign layer. The material is very hydrophobic in the starting state and very quickly regains hydrophobicity after the imposed loss by means of the plasma and therefore exhibits an excellent recovery effect.

The second Example exhibits virtually the same hydrophobicity transfer effect as Example 1, but the surface hydrophobicity is not as good. After the loss thereof (by means of plasma), hydrophobicity is restored, but less quickly than in Example 1. The third Example exhibits a very good hydrophobicity transfer effect which is established rapidly.

3) The further Comparison Examples show that the recovery effect cannot be influenced by the individual additives alone. In all cases the hydrophobicity is not regained after imposed loss.

Application Example 3 (comparative cracking test):

Shaped steel articles are encapsulated either with casting resin system 3 according to the invention (see IE3) or with an unmodified casting resin system and the resin is cured. Then in a cycle having defined time units, the shaped articles are exposed to specific periods at steadily increasing temperatures and after each period are examined for any crack formation which may have occurred.

Preparation of the unmodified comparison mixture (analogously to Comparison Example 1): In a mixing apparatus having a blade stirring mechanism, 1000 g of epoxy resin 1, 900 g of hardener 1, 5.0 g of BDMA and 3780 g of quartz powder W 12 EST are mixed together in the course of 30 min at 60°C using a mixer having a blade stirring mechanism and then briefly degassed at about 10 mbar.

The shaped articles are subjected to the cracking test described below:

Cracking test

In an aluminium casting mould having depressions, batches of 20 steel test specimens of the casting resin systems to be compared are produced by means of the APG (Automatic Pressure Gelation) process. In that process, the liquid resin mixture is heated to 40-60°C and stirred in vacuo. The resin mixture is then injected into the casting mould (which has been heated to 140-150°C) under a slight pressure of 2-5 bar. While the pressure is maintained, the resin system is gelled in the casting mould within a period of a few minutes, each test specimen produced thus having an enclosed steel body. After gelation, the test specimens are removed from the mould and post-cured for 10 hours at 140°C. The test specimens are then exposed to a specific temperature profile. The temperature profile consists of a series of cycles having an upper temperature limit of 25°C, at which each specimen is also examined for any crack formation, whereas the lower temperature limit becomes lower with every cycle. In each cycle, the specimens are rapidly brought to the respective lower or upper temperature limit, and maintained at that particular limit temperature for a few hours. The cycle in which a specimen exhibits the first cracks is noted. The average cracking temperature of a resin system is determined from the frequency distribution of crack formation per cycle.

The results of the comparative cracking test of Application Example 3 are listed in the following Table:

Temperature [°C]	IE (20 specimens)	CE (20 specimens)
/ duration [hours]	cracked specimens	cracked specimens
25 / 8	0	0
-10 / 8 + 25 / 8	0	17
-20 / 16 + 25 / 8	1	2
-40 / 16 + 25 / 8	4	1 (20)
- 60 /16 + 25 / 8	7	-
- 80 / 16 + 25 / 8	5	-
- 100 / 16 + 25 / 8	3 (20)	-

In the case of the comparison system, the average cracking temperature is calculated to be -12°C, whereas it is -66°C for the system according to the invention. The systems according

to the invention, in addition to their hydrophobic behaviour, therefore surprisingly also exhibit an appreciably better temperature change behaviour in comparison with an unmodified system.

Patent claims

- 1. A composition comprising
- (a) an epoxy resin,
- (b) an OH-terminated polysiloxane,
- (c) a cyclic polysiloxane and
- (d) a non-ionic, fluoroaliphatic surface-active reagent.
- 2. A composition according to claim 1, comprising, based on the total composition (a), (b), (c) and (d), from 77.0 to 97.99 % by weight component (a), from 1.0 to 10.0 % by weight component (b), from 1.0 to 10.0 % by weight component (c) and from 0.01 to 3.0 % by weight component (d), the sum of components (a), (b), (c) and (d) being 100 % by weight.
- 3. A composition according to claim 1, comprising as component (a) a cycloaliphatic epoxy resin or an epoxidation product of a natural unsaturated oil or a derivative thereof.
- 4. A composition according to claim 1, comprising as component (a) hexahydrophthalic acid diglycidyl ester and 3,4-epoxycyclohexylmethyl 3',4'-epoxycyclohexanecarboxylate as well as epoxidised soybean oil or epoxidised linseed oil.
- 5. A composition according to claim 1, comprising as component (b) a polysiloxane of formula I

$$HO = \begin{bmatrix} R_1 \\ I \\ SI - O \end{bmatrix} H \qquad (I)$$

wherein R_1 and R_2 are each independently of the other C_1 - C_{18} alkyl, C_5 - C_{14} aryl or C_6 - C_{24} -aralkyl and n is an average value of from 3 to 60.

- 6. A composition according to claim 5, comprising as component (b) a polysiloxane of formula I wherein R_1 and R_2 are each independently of the other methyl, ethyl or phenyl.
- 7. A composition according to claim 5, comprising as component (b) a polysiloxane of formula I wherein R_1 and R_2 are methyl.
- 8. A composition according to claim 1, comprising as component (c) a cyclic polysiloxane of formula II

$$\begin{array}{c|c}
R_1 \\
Si - O \\
R_2
\end{array}$$
(II),

wherein R_1 and R_2 are each independently of the other C_1 - C_{18} alkyl, C_5 - C_{14} aryl or C_6 - C_{24} -aralkyl and m is an integer from 3 to 12.

- 9. A composition according to claim 8, comprising as component (c) a cyclic polysiloxane of formula II wherein R_1 and R_2 are each independently of the other methyl, ethyl or phenyl and m is an integer from 3 to 8.
- 10. A composition according to claim 8, comprising as component (c) octamethylcyclotetrasiloxane, decamethylcyclopentasiloxane, dodecamethylcyclohexasiloxane or a hydrolysate of dimethyldichlorosilane.
- 11. A composition according to claim 11, comprising as component (d) a non-ionic, fluoroaliphatic surface-active reagent of formula

$$R_f$$
 -COO-(CH₂CH₂O)_m-R_H (III)

or

$$R_f - (CH_2CH_2O)_m - R_H$$
 (IV),

wherein m = 1 to 200, R_f is a linear or branched perfluorinated alkyl having from 2 to 22 carbon atoms and R = H, C_1 - C_6 alkyl or R_f .

- 12. A composition according to claim 11, comprising as component (d) compounds of formula (III) or (IV) in which the molecular mass according to the theoretical empirical formula is from 300 to 8000.
- 13. A composition according to claim 12, comprising as component (d) 1,1,2,2-tetrahydro-perfluorooctanol or R_f -COO-(CH_2CH_2O)_m-R, wherein R_f is a linear perfluorinated alkyl having from 16 to 18 carbon atoms, m = 110-130 and R = H.
- 14. A composition according to claim 1, additionally comprising as further component (g) emulsifiers or thickeners.
- 15. A composition according to claim 14, comprising as component (g) from 0.01 to 3.5 % by weight, based on the sum of components (a) to (d), highly dispersed, hydrophilic, untreated silicic acid.
- A composition according to claim 1 or 14 or 15, additionally comprising a curing agent.
- 17. A composition according to claim 16, wherein the curing agent is selected from a polycarboxylic anhydride (e) or from a polycarboxylic anhydride (e) together with an accelerator (f).
- 18. A composition according to claim 16, wherein the curing agent is an initiator system for the cationic polymerisation.
- 19. A composition according to any one of claims 1 or 14 to 18, which comprises fillers.
- 20. A composition according to claim 19, which comprises as filler quartz powder, silanised quartz powder, aluminium hydroxide or aluminium oxide.
- 21. A crosslinked product obtainable by curing a composition according to any one of claims 1 to 20.

22. An electrical insulating material according to claim 21.

Abstract of the Disclosure

A composition comprising

- (a) an epoxy resin,
- (b) an OH-terminated polysiloxane,
- (c) a cyclic polysiloxane and
- (d) a non-ionic, fluoroaliphatic surface-active reagent, exhibits excellent hydrophobicity properties and can be used as electrical insulating material.